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## **Analysis of Hair Samples for Sympathomimetic Amines**

#### 1 Introduction

Sympathomimetic amines (SMAs) are generally a class of synthetic phenethylamine-derived drugs often generically referred to as "amphetamines". Almost all of these compounds show some degree of stimulant effects, but a wide variety of additional structure-dependent pharmacological effects can be seen in various compounds. These include pure stimulants (amphetamine and methamphetamine), decongestants (phenylpropanolamine and pseudoephedrine), anorexics (phentermine and fenfluramine), and hallucinogens (mescaline, one of the few relevant naturally occurring SMAs).

## 2 Scope

This procedure allows for screening, confirmation and quantitation of the following SMAs in hair samples: amphetamine, methamphetamine, ephedrine / pseudoephedrine, methylenedioxyethylamphetamine (MDEA), and methylenedioxymethamphetamine (MDMA). It also allows for the qualitative analysis of hair samples for methylenedioxyamphetamine (MDA). This document applies to Chemistry Unit case working personnel who perform toxicology analyses.

## 3 Principle

Hair samples may be qualitatively or quantitatively assayed for SMAs. After washing, specimens are dried and cryoground into a powder. The resulting hair powder is mixed with an internal standard (normally a mixture of six deuterated SMAs) and extracted in methanol overnight. The methanol extracts are then taken to dryness and reconstituted in water. The aqueous extracts are adjusted to a basic pH, and extracted with hexane. The hexane is removed, acidified to prevent evaporation of volatile SMAs, and taken to dryness. The resulting residue is reconstituted in 10/90 methanol/water and analyzed by liquid chromatography with high resolution mass spectrometry (LC-FTMS). The extraction procedure is derived from work by Sadeghipour and Veuthey. The chromatographic and mass spectral procedures and parameters were developed in-house.

## 4 Specimens

This procedure uses 50 mg of hair if specimens are analyzed in duplicate.

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## 5 Equipment/Materials/Reagents

- a. 16x100 mm screw-top tubes with Teflon-lined caps
- b. 12x75 mm culture tubes with polypropylene snap-tops
- c. Acetonitrile (Optima grade or better)
- d. Formic Acid (Puriss grade or better)
- e. Hexane (UV grade or better)
- f. Hydrochloric acid (ACS grade or better)
- g. Methanol (Optima grade or better)
- h. Sodium hydroxide (ACS grade or better)
- i. Water (Deionized and Optima or better grade)
- j. 4% Sodium hydroxide Dissolve 2 g sodium hydroxide in 50 mL deionized water. Store in plastic at room temperature. Stable for at least 6 months.
- Methanol:Hydrochloric Acid (4:1 v:v)
   Mix 20 mL methanol with 5 mL hydrochloric acid. Store in glass at room temperature.
   Stable for at least 1 month.
- Methanol: Water (10:90 v:v)
   Mix 5 mL methanol with 45 mL water (both Optima grade or better). Store in glass at room temperature. Stable for at least 1 year.
- m. 0.1% Formic acid in acetonitrile
   Obtain 500 mL acetonitrile and mix with 0.5 mL formic acid. Store in glass at room temperature. Stable for 2 months.
- n. 0.1% Formic acid in water Vacuum filter 500 mL water (Optima grade or better) through a 5  $\mu$ m PTFE membrane and mix with 0.5 mL formic acid. Store in glass at room temperature. Stable for 2 months.
- o. Vortex mixer, Rotator, Centrifuge, Heating Block and Cryogrinder
- p. Evaporator with nitrogen

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- q. Routine laboratory supplies, including disposable pipettes, wooden sticks, test tube racks, graduated cylinders, etc.
- r. Liquid Chromatograph-Orbitrap Mass Spectrometer
- s. HPLC Column (Alltech Alltima C18, 2.1 x 150 mm, 5 μm dp, with a 2.1 x 7.5 mm guard column; or equivalent)
- t. Methylene chloride (HPLC grade)
- u. Eppendorf Thermomixer
- v. Ultrafree-CL centrifuge filters (0.45 µm PVDF)

#### 6 Standards and Controls

Separate sources for calibration and control material are required (see TOX101).

- a. d<sub>3</sub>-Ephedrine Stock Solution (100 μg/mL):
   A methanol solution purchased from Cerilliant or other approved vendor. Stability and storage conditions are determined by the manufacturer.
- b. d<sub>5</sub>-Amphetamine Stock Solution (100 μg/mL):
   A methanol solution purchased from Cerilliant or other approved vendor. Stability and storage conditions are determined by the manufacturer.
- c. d<sub>5</sub>-Methamphetamine Stock Solution (100  $\mu g/mL$ ): A methanol solution purchased from Cerilliant or other approved vendor. Stability and storage conditions are determined by the manufacturer.
- d. d5-MDA Stock Solution (100 μg/mL):
   A methanol solution purchased from Cerilliant or other approved vendor. Stability and storage conditions are determined by the manufacturer.
- e. d<sub>5</sub>-MDMA Stock Solution (100 μg/mL):
   A methanol solution purchased from Cerilliant or other approved vendor. Stability and storage conditions are determined by the manufacturer.
- f. d<sub>5</sub>-MDEA Stock Solution (100 μg/mL):
   A methanol solution purchased from Cerilliant or other approved vendor. Stability and storage conditions are determined by the manufacturer.

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- g. Internal Standard Working Solution (2 μg/mL of each component): Combine 0.5 mL each of the d<sub>3</sub>-ephedrine, d<sub>5</sub>-amphetamine, d<sub>5</sub>-methamphetamine, d<sub>5</sub>-MDA, d<sub>5</sub>-MDMA, and d<sub>5</sub>-MDEA stock solutions in a 25 mL volumetric flask. Add 2 mL methanol and bring to the mark with water (both Optima grade or better). Store in glass at <0°C. Stable for at least 2 years.
- h. Hair Internal Standard Working Solution (25 ng/mL of each component): Add 625  $\mu$ L of the Internal Standard Working Solution (2  $\mu$ g/mL) to a 50 mL volumetric flask and bring to the mark with Optima grade methanol. Store in glass at <0°C. Stable for at least 2 years.
- i. Ephedrine Stock Solution (1 mg/mL):
   A methanol solution purchased from Cerilliant or other approved vendor. Stability and storage conditions are determined by the manufacturer.
- j. MBDB (N-methylbenzodioxazolylbutanamine, N-methyl-1-(3,4-methylenedioxyphenyl)-2-butanamine) Stock Solution (1 mg/mL):
   A methanol solution purchased from Cerilliant or other approved vendor. Stability and storage conditions are determined by the manufacturer.
- k. Amine Mixture-6 (250 µg/mL each component):
   A methanol solution containing amphetamine, methamphetamine, phentermine, MDA, MDMA, and MDEA purchased from Cerilliant or another approved vendor. Stability and storage conditions are determined by the manufacturer.
- 1. Column Performance Evaluation Mix (1  $\mu$ g/mL each component): Combine 25  $\mu$ L each of the MBDB and ephedrine stock solutions with 100  $\mu$ L of the Amine Mixture-6 in a 25 mL volumetric flask. Add 2.4 mL methanol and bring to the mark with water (both Optima grade or better). Stable for at least 2 years. A 10  $\mu$ L portion of this solution is analyzed before each day's samples, in order to confirm acceptable instrument performance.
- m. Amphetamine Stock Solution (1 mg/mL):
  A methanol solution purchased from Cerilliant or other approved vendor. Stability and storage conditions are determined by the manufacturer.
- n. Methamphetamine Stock Solution (1 mg/mL):
   A methanol solution purchased from Cerilliant or other approved vendor. Stability and storage conditions are determined by the manufacturer.
- o. MDA Stock Solution (1 mg/mL):
   A methanol solution purchased from Cerilliant or other approved vendor. Stability and storage conditions are determined by the manufacturer.

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p. MDMA Stock Solution (1 mg/mL):

A methanol solution purchased from Cerilliant or other approved vendor. Stability and storage conditions are determined by the manufacturer.

q. MDEA Stock Solution (1 mg/mL):

A methanol solution purchased from Cerilliant or other approved vendor. Stability and storage conditions are determined by the manufacturer.

- r. Control Working Solution (1 μg/mL each component):
  - Mix 50  $\mu$ L each of the ephedrine, amphetamine, methamphetamine, MDA, MDMA, and MDEA stock solutions in a 50 mL volumetric flask. Add 7 mL methanol and bring to the mark with water (both Optima grade or better). Store in glass at <0°C. Stable for at least 1 year.
- s. Hair Control Working Solution (125 ng/mL each component): Add 1.25 mL of the Control Working Solution (1  $\mu$ g/mL) to a 10 mL volumetric flask and bring to the mark with Optima grade methanol. Store in glass at <0°C. Stable for at least 1 year.
- t. Calibration Working Solution #1 (5  $\mu$ g/mL each component): Mix 250  $\mu$ L each of the ephedrine, amphetamine, methamphetamine, MDA, MDMA, and MDEA stock solutions in a 50 mL volumetric flask. Add 8.5 mL methanol and bring to the mark with water (both Optima grade or better). Store in glass at <0°C. Stable for at least 1 year.
- u. Calibration Working Solution #2 (0.5  $\mu$ g/mL each component): Mix 25  $\mu$ L each of the ephedrine, amphetamine, methamphetamine, MDA, MDMA, and MDEA stock solutions in a 50 mL volumetric flask. Add 9.9 mL methanol and bring to the mark with water (both Optima grade or better). Store in glass at <0°C. Stable for at least 1 year.
- v. Hair Calibration Working Solution #3 (125 ng/mL each component): Add 625  $\mu$ L of the Calibration Working Solution #1 (5  $\mu$ g/mL) to a 25 mL volumetric flask and bring to the mark with Optima grade methanol. Store in glass at <0°C. Stable for at least 1 year.
- w. Hair Calibration Working Solution #4 (12.5 ng/mL each component): Add 625  $\mu$ L of the Calibration Working Solution #2 (0.5  $\mu$ g/mL) to a 25 mL volumetric flask and bring to the mark with Optima grade methanol. Store in glass at <0°C. Stable for at least 1 year.
- x. Negative Control Hair:

Purchased from Diagnostics Products Corporation, UTAK Laboratories, Inc., Cliniqa, or prepared in-house from an appropriate blank specimen. Hair will be stored at room

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temperature, and does not expire. A Negative Control hair sample will be extracted and analyzed with every assay.

## y. Positive Control Hair:

This is normally prepared in-house as per the *Quality Control for Toxicology Examinations* technical procedure (TOX101), but may be purchased from an appropriate vendor as needed. When prepared in house, it will be made fresh. Normally prepared at concentrations of 50 and 500 pg/mg by adding 10 and 100  $\mu$ L of the Hair Control Working Solution to 25 mg of Negative Control Hair. Other levels may be used as circumstances dictate.

This procedure may be used quantitatively via construction of a multi-point calibration curve for the analyte(s) of interest following the *Quality Control for Toxicology Examinations* technical procedure (TOX101). Table 1 shows typical calibrators and preparation instructions for hair calibrators.

Table 1: Hair Calibrator Preparation

Cal Level (pg/mg)	Hair Amount (mg)	Hair Calibrator Working Solution #3 Volume (μL)	Hair Calibrator Working Solution #4 Volume (μL)
25	25	0	50
50	25	0	100
75	25	0	150
175	25	35	0
300	25	60	0
500	25	100	0
750	25	150	0
1000	25	200	0

## 7 Sampling

Representative portions of the specimens are obtained. See TOX101 for further details.

#### 8 Procedure

Appendix 1 contains an abbreviated version of this procedure. This form may be used at the bench by the authorized individual performing the procedure.

## 8.1 Preparation of Hair Samples:

Visually inspect hair and record observations.

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- b. If segmental analysis is required, cut a portion of the hair sample into 2-cm segments.
- c. Accurately weigh 25-100 mg of each hair sample into a properly labeled test tube (to the nearest 0.1 mg).
- d. Wash each hair sample with 1.5 mL methanol by vortexing for approximately 1 minute. Discard this wash.
- e. Wash each hair sample with 1.5 mL methylene chloride by vortexing the sample for approximately 1 minute. Discard this wash.
- f. Wash each hair sample with 1.5 mL methanol by vortexing for approximately 1 minute. Save this final wash for later analysis, if necessary. Control washes need not be saved.
- g. Dry hair samples in a Thermomixer/heating block at approximately 40°C to evaporate any remaining solvent.
- h. Cryogrind dry hair samples in the freezer mill using the settings in Section 9.3 of this procedure.
- i. Accurately weigh 25 mg of hair powder to a small vial (to the nearest 0.1 mg). Samples will be prepared in duplicate if specimen size allows. Smaller amounts may be weighed to account for high concentrations of analyte and/or limited specimen amount.
- j. Add 1.5 mL methanol, and 50  $\mu$ L Hair Internal Standard Working Solution (25 ng/mL) to each vial.
- k. Extract overnight (at least 12 hours) with stirring at 37°C in a Thermomixer.
- 1. Filter the methanol extract using an Ultrafree-CL 0.45 μm centrifuge filter by spinning at 3000 rpm for 5 minutes. Discard the hair.
- m. Add 0.1 mL of 4:1 methanol:hydrochloric acid to filtrate and vortex briefly.
- n. Evaporate to dryness under a gentle stream of nitrogen at approximately 40°C.
- o. Reconstitute each sample in 0.5 mL deionized water by vortexing for at least 10 seconds.

#### 8.2 For Hair Extracts:

- a. To a properly labeled 16x100 mm screw-top tube add 0.5 mL of water based hair extract.
- b. Add 0.2 mL of 4% sodium hydroxide to each sample and vortex briefly.

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- c. Add 2 mL of hexane to each tube and extract for 20 minutes on a rotator. Centrifuge 10 minutes at a minimum of 3000 rpm. Use a wooden stick to break up any emulsions that develop.
- d. Transfer organic (top) layer to a 12x75 mm culture tube.
- e. Add 0.1 mL of 4:1 methanol:hydrochloric acid and vortex briefly.
- f. Evaporate the hexane to dryness under a gentle stream of nitrogen at approximately 40°C.
- g. Reconstitute the dried residue in 0.1 mL of 10:90 methanol:water.
- h. Analyze by LC-FTMS using the conditions given below (Sections 9.1 and 9.2).

## 8.3 For Wash Samples:

- a. For samples in which an SMA is identified above the LLOQ of the method, add 25  $\mu$ L Hair Internal Standard Working Solution (25 ng/mL) to each wash.
- b. Add 0.1 mL of 4:1 methanol:hydrochloric acid and vortex briefly.
- c. Evaporate to dryness under a gentle stream of nitrogen at approximately 40°C.
- d. Reconstitute each sample in 0.5 mL deionized water by vortexing for at least 10 seconds.
- e. Add 0.2 mL of 4% sodium hydroxide to each sample and vortex briefly.
- f. Add 2 mL of hexane to each tube and extract for 20 minutes on a rotator. Centrifuge 10 minutes at a minimum of 3000 rpm. Use a wooden stick to break up any emulsions that develop.
- g. Transfer organic (top) layer to a 12x75 mm culture tube.
- h. Add 0.1 mL of 4:1 methanol:hydrochloric acid and vortex briefly.
- i. Evaporate the hexane to dryness under a gentle stream of nitrogen at approximately 40°C.
- j. Reconstitute the dried residue in 0.1 mL of 10:90 methanol:water.
- k. Analyze by LC-FTMS using the conditions given below (Sections 9.1 and 9.2).

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## 9 Instrumental Conditions

Appendix 2 contains a checklist of method parameters that should be used to verify proper instrumental conditions prior to analysis of case samples.

## 9.1 Liquid Chromatograph Parameters (Shimadzu Prominence, or equivalent)

Mobile Phase Compositions	Flow Parameters		Column Parameters		
B: 0.1% formic acid in	total flow 0.3 mL/		L/min	type	C18
acetonitrile	time (min)	%B	%C	length	15 cm
C: 0.1% formic acid in water	0	7.5	92.5	internal diameter	2.1 mm
	5	7.5	92.5	particle size	5 μm
	20	60	40	temperature	40°C
	23	60	40	guard length	7.5 mm
	28	7.5	92.5	guard ID	2.1 mm
	32	7.5	92.5		
	total time	32 mi	n		

## 9.2 Mass Spectrometer Parameters Using FTMS (Thermo Orbitrap, or equivalent)

Source Parameters				
Mode: Electrospray		Spray Voltage: +5 kV	Capillary Temperature:	
			250°C	
Sheath Gas: 25 (arb units)		Aux Gas: 10 (arb units)	Sweep Gas: 0 (arb units)	
All other source parameters are set through the tuning process. See the appropriate IOSS				
technical procedure for details.				
Scan Range	100-350 m/z			
Resolution	30000			

## 9.3 Cryogrinder (Freezer/Mill) Parameters

Cycles	1
Precool	Auto
Run time	6.5 min
Rate	25 hz

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#### 10 Decision Criteria

The following criteria are used as guidelines in determining the acceptability of the data produced in this assay. In general, compound identification should be based on a comparison of the chromatography and mass spectrometry for the analyte peak of interest with data from a contemporaneously analyzed reference standard, calibrator, or extracted Positive Control.

### 10.1 Chromatography

The peak of interest should show good chromatographic fidelity, with reasonable peak shape, width, and resolution. In order to be determined acceptable, a chromatographic peak in an unknown sample should compare favorably to a chromatographic peak of the same analyte in a known sample analyzed on the same system in the same or subsequent analytical runs. Additionally, the following two criteria should be met.

#### 10.1.1 Retention Time

The retention time of the peak should be within  $\pm$  5% of the retention time (relative or absolute, as appropriate) obtained from injection of a reference standard, calibrator, or Positive Control.

## 10.1.2 Signal-to-Noise

To justify the existence of a peak, its baseline signal to peak-to-peak noise ratio should exceed 3. Further, the baseline signal for the peak of interest should be at least 10 fold greater than that for any observed peak at similar retention time in a Negative Control or blank injected just prior to the sample.

#### 10.2 Mass Spectrometry

The M+1 for the compound of interest should agree with the theoretical exact mass within 0.003 amu. See Table 2 below for theoretical exact masses.

Table 2: Theoretical Exact Masses (M+1)

Compound Name	Exact Mass (M+1)
Amphetamine	136.112
Methamphetamine	150.128
Ephedrine	166.123
MDA	180.102
MDMA	194.118
MDEA	208.133

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#### 10.3 Wash Decision Criteria

If the final wash contains greater than one tenth the amount of an SMA in a given hair sample, the sample's exterior is considered to be possibly contaminated, and will be reported as such.

#### 11 Calculations

See the *Quality Control for Toxicology Examinations* technical procedure (TOX101) for acceptable practices in calculating quantitative results. Equal weighting is used for quantitated analytes.

For wash calculations, the total amount of an SMA may be calculated against a new curve, or against the hair curve for that sample.

Example calculations for wash decision criteria:

Assume 25.0 mg of sample Q1 are washed per the SOP, and that the sample is determined to contain 500 pg methamphetamine per milliliter of hair. Then, the hair sample contained 12500 pg total of methamphetamine (25 x 500). The final wash must contain more than 1250 pg methamphetamine (12500/10) for the Q1 sample to be reported as possibly contaminated.

## 12 Measurement Uncertainty

The critical sources of measurement uncertainty in this procedure include:

- historical random uncertainty of repeated measurements
- accuracy of the balance used to deliver the sample
- accuracy of the pipette used to deliver the calibrators
- uncertainty in the concentration of the calibration standards
- precision of the delivery of internal standard

When quantitative results are included in an FBI Laboratory report, the measurement uncertainty will be estimated and reported following the *Chemistry Unit Procedures for Estimating Measurement Uncertainty technical* procedure (CUQA 13). Information used to derive uncertainty measurements will be tracked in an electronic database.

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### 13 Limitations

a. Method Performance Parameters:

Compound	LOD (pg/mg)	LLOQ (pg/mg)	Linear Range (pg/mg)	Accuracy (% bias at low and high controls)	Precision (% intermediate at low and high controls)
Amphetamine	21	25	25-1000	+3.81, +22.61	17.63, 17.54
Methamphetamine	14	25	25-1000	+1.72, +6.48	7.77, 8.79
(pseudo)Ephedrine	25	25	25-1000	-0.52, +9.99	11.76, 14.85
MDMA	14	25	25-1000	+5.66, +10.78	7.12, 5.39
MDEA	8	25	25-1000	-2.74, +5.33	9.14, 5.32
MDA	25			N/A; Qual Only	

b. Interferences: None known.

## 14 Safety

Take standard precautions for the handling of chemicals and biological materials. Refer to the *FBI Laboratory Safety Manual* for guidance.

#### 15 References

Sadeghipour, F. and Veuthey, J., Journal of Chromatography A, v. 787 (1997), pp. 137-143

Baselt, R.C., *Disposition of Toxic Drugs and Chemicals in Man*, 7th ed., Biomedical Publications: Foster City, California, 2004.

Quality Control for Toxicology Examinations (TOX101); FBI Laboratory Chemistry Unit – Toxicology SOP Manual.

Chemistry Unit Procedures for Estimating Measurement Uncertainty (CUQA 13); FBI Laboratory Chemistry Unit Quality Assurance and Operations Manual.

Guidelines for Comparison of Mass Spectra (Tox 104); FBI Laboratory Chemistry Unit – Toxicology SOP Manual.

FBI Laboratory Chemistry Unit – Instrument Operation and Support SOP Manual.

FBI Laboratory Safety Manual.

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0	03/08/2012	n/a	New document. SMA hair analysis was pulled out of
			Tox 420 into this free standing document.
1	09/01/2021	(7)	Removed calibration section and renumbered
		7	Updated sampling language
		5, 15	Removed references to TOX103
		Multiple	Removed "subunit"
		1	Removed excess language
		2	Updated scope language
		3	Clarified that qualitative is also an option
		5-m	Removed pre-filtering of mobile phase
		6 Clarified separate sources for calibrator and co	
		8 Updated language to "authorized individual",	
		subdivided section for clarity	
		8-g	Added Thermomixer as a heat source
		11	Added calibration curve weighting
		12, 15	Updated section and SOP title
		6, 11, 15	Updated TOX101 title
		9.3	Updated cryogrinder parameters
		5-u, 8-j, k, 1	Removed magnetic stirbar references, replaced with
			Thermomixer
		Multiple	Replaced "standard operating procedure" with
			"technical procedure". Corrected minor
			typographical/formatting errors on bench sheets.

# Approval Redact - Signatures on File

Chemistry Unit Chief: Date: 08/31/2021

Toxicology Technical

Leader: Date: 08/31/2021

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# Appendix 1: Abbreviated version of the SMA procedure for bench use. (Page 1 of 2)

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# Appendix 1: Abbreviated version of the SMA procedure for bench use. (Page 2 of 2)

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# Appendix 2: Instrumentation parameters checklist for the SMA procedure.

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